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		Application Number	10/806,899
		Filing Date	3/23/2004
		First Named Inventor	Petrou et al.
		Art Unit	1614 1634
		Examiner Name	Stephen Thomas Kapushoc
Sheet	1	of	2
		Attorney Docket Number	

U.S. PATENT DOCUMENTS

FOREIGN PATENT DOCUMENTS

Examiner Initials*	Cite No. ¹	Foreign Patent Document	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	T ⁶
		Country Code ³ - Number ⁴ - Kind Code ⁵ (if known)				
/STK/	3	WO 2005/014863	02-17-2005	Bionomics Limited		
/STK/	4	WO 2004/085674	10-07-2004	Bionomics Limited		
/STK/	5	WO 2002/050096	06-27-2002	Bionomics Limited		
/STK/	6	WO 2002/006521	01-24-2002	Bionomics Limited		

Examiner Signature	/Stephen Kapushoc/	Date Considered	08/04/2008
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Substitute for form 1449B/PTO				Complete if Known	
				<i>Application Number</i>	10/806,899
				<i>Filing Date</i>	3/23/2004
				<i>First Named Inventor</i>	Petrou et al.
				<i>Art Unit</i>	1614 1634
				<i>Examiner Name</i>	Stephen Thomas Kapushoc
Sheet	2	of	2	Attorney Docket Number	1386/19

NON PATENT LITERATURE DOCUMENTS						
Examiner Initials*	Cite No.	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.				T ²
/STK/	7	Chou et al., "The lack of association between febrile convulsions and polymorphisms in SCN1A," Epilepsy Research, Vol. 54, pgs. 53-57 (2003).				
/STK/	8	Fujiwara et al., "Mutations of sodium channel a subunit type 1 (SCN1A) in intractable childhood epilepsies with frequent generalized tonic-clonic seizures," Brain, Vol. 126, pgs. 531-546 (2003).				
/STK/	9	Hirschhorn et al., "A comprehensive review of genetic association studies," Genetics in Medicine, Vol. 4, No. 2, pgs. 45-61 (2002).				
/STK/	10	Notification Concerning Transmittal of Copy of International Preliminary Report on Patentability for International Application No. PCT/AU2006/000841 dated January 3, 2008.				
/STK	11	Official Action for U.S. Patent Application Serial No. 10/482,834 dated August 2, 2007.				
/STK/	12	Official Action for U.S. Patent Application Serial No. 10/482,834 dated April 4, 2008.				
/STK	13	Official Action for U.S. Patent Application Serial No. 11/262,647 dated February 15, 2008.				
/STK	14	Ohmori et al., "Significant correlation of the SCN1A mutations and severe myoclonic epilepsy in infancy," Biochemical and Biophysical Research Communications, Vol. 295, pgs. 17-23 (2002).				
/STK/	15	Stafstrom et al., "Epilepsy Genes: The Link Between Molecular Dysfunction and Pathophysiology," Mental Retardation and Developmental Disabilities Research Reviews, Vol. 6, pgs. 281-292 (2000).				
/STK/	16	Supplementary European Search Report corresponding to Australian Patent No. AU0200910 dated February 17, 2005.				

Examiner Signature	/Stephen Kapushoc/	Date Considered	08/04/2008
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